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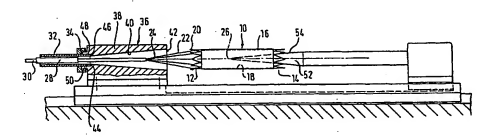
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(54) Title: METHOD OF, AND DEVICE FOR, INSTALLING A STENT IN A SLEEVE



(57) Abstract: A device for compressing and pulling into the sleeve of a stent delivery system, in a pull direction along a long axis, a self-expanding stent which has a long axis which is coaxial with that of the delivery system. The pulling device includes a funnel to lead the stent into the sleeve, a pull structure to be received with the sleeve, a clamp to restrain the sleeve from following the stent and means to release the pull structure from the stent.

METHOD OF, AND DEVICE FOR, INSTALLING A STENT IN A SLEEVE

Field of the Invention

This invention relates to devices and methods for installing a self-expanding stent into a confining sleeve at the distal end of a stent delivery system.

Background

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Among the types of stents that may be placed within a human vessel are those characterized as "self-expanding" stents which are confinable, in a small diameter, in the distal end of a catheter-like delivery device in readiness for advancement to and release within a selected site in the vessel. Such stents may be made of a shaped memory alloy, such as a nickel-titanium alloy. For example, such stents may be made from a cylinder of the alloy with a diameter corresponding to, but usually somewhat less than, the diameter which the stent has in its compressed delivery configuration, that is, when it is confined within a sheath of a stent delivery system, for introduction into the body of the patient, for placement at the desired site within the body. When the stent is placed at the desired location in the vessel, the confining sheath is withdrawn proximally, allowing the compressed stent to expand, from memory, to an expanded configuration in which it supports the wall of the bodily lumen in which it is sited.

In the production process, the starting cylinder may be laser-cut to create a matrix of through openings, and then expanded to expanded diameter. It is then heated to give it a "memory" of this configuration. After that, it is necessary to compress the stent sufficient to introduce it into the sleeve or sheath of the delivery system. The compression and loading of the stent into the sleeve may be performed manually and may involve using a fixture, such as a conical funnel through which the stent is urged to progressively compress it to a smaller diameter in which it can be installed into the sleeve. The stent may be covered by an appropriate graft material, such as expanded polytetrafluoroethylene (ePTFE), defining a stent-graft. Alternately, the graft may be a bare, uncovered stent. Where the compression and loading of the stent into the sleeve involves a number of manual steps, there is significant risk of variation in the manner in which the stent engages and is seated within the sleeve.

One object of the present invention is to provide a stent compression process and apparatus which will allow the step of installing the stent in its delivery sheath or sleeve to be conducted in a more automated way, which should facilitate the objectives of strict quality control, predictability, and further process improvement.

- Also among the objects of the invention are to provide a process and apparatus for controlling the degree of longitudinal force applied to the stent; to accurately and repeatedly position the stent at a predetermined distance from the distal end of the delivery sheath; to support the stent from within during its compression.
- In EP-A2-0 657 147, C. R. Bard, Inc. proposes to install a stent graft in a sleeve by a process which involves pushing a leading end of the stent graft into a conical confining funnel, until the leading end of the stent, at the tip of the funnel, at its compressed configuration, engages with radially extending spokes of an implant retention device within the sleeve of the delivery system. The spokes act within the sleeve as an anchor, to prevent undesirable axial movements of the stent graft within the sheath/sleeve, or in deployment of the stent graft from the sheath. As the implant expands on deployment, it moves radially outwardly away from the spokes of the anchor device, thereby automatically releasing the anchor device for withdrawal from the blood vessel.

US-A-5,591,222, Susawa et al includes a disclosure of the use of a wire to draw a polymeric stent into a confining tube, as part of the process of placing the stent over an angioplasty balloon.

US-A-5,649,950 Bourne et al discloses use of conical-shaped lumen for collapsing a prosthetic occluder into a narrow or slender configuration advanceable through the lumen of an introducer sheath. The system is described as a front-end loader and contrasted with the Rashkind prior art rear-end loading delivery system. The prosthesis is held on a locking wire, but is drawn by a plurality of suture lines down the conical confinement surface, in a direction opposite to that in which the wire extends away from the prosthesis.

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US-A-5749921, Lenker et al discloses loading a Nitinol® stent into a tube by pulling it down a funnel using a plurality of pulling filaments, each looped around a different one of the apices of the stent mesh at its leading end. During deployment of the stent at a site of surgery, the threads can be used once again to pull the stent back into the tube, should this be required. To release the stent, one end of each loop is cut and the other end is used to pull the thread through the stent apex until the cut end passes through the apex

WO 97/17021 Dereume discloses a device to recover into a lumen a stent previously placed in a bodily lumen. The device deploys a ring of hooks to engage one end of the stent and pull it down a flared end of a lumen of the elongate recovery device. See also WO 2000/02615 and CA 2213291 (Gianotti).

The manual stent installation process for a nickel-titanium stent also may involve cooling the stent to below room temperature, in a water bath. While this is acceptable for an uncoated metal stent, which can be subsequently sterilized totally, it is less attractive for a coated stent or a stent graft, because of the potential for the prosthesis to carry bacteria and pyrogens, originating in the water bath. Thus, it is another object of the present invention to find a method to install a self-expanding metal stent within the sleeve of a delivery system, in a gaseous rather than liquid environment, but which also offers the potential to control the temperature of the stent being installed, especially to bring it to temperatures below room temperature.

Summary of the Invention

In one aspect of the present invention there is provided a device for installing a stent in the lumen of a sleeve: the stent having luminal and abluminal surfaces, a leading end and a trailing end and having a capacity to expand radially from a compressed configuration to an expanded configuration; the sleeve having a long axis, a lumen, a proximal receiving end and a distal end near which the stent is to be installed; the device comprising: an elongate puller, to be received in the sleeve and having a leading end and a trailing end; an adaptor, for coupling the trailing end of the puller to the leading end of the stent when the stent is in its expanded configuration and for decoupling the puller from the stent when the stent is in its compressed configuration; a

gripper, to grip the receiving end of the sleeve; a constrictor, to compress and guide the leading end of the stent into the receiving end of the sleeve; a base; a puller holder, to transfer stresses between the puller and the base; a gripper holder, to transfer stresses between the gripper and the base; and a translator, to increase the distance between the puller and the gripper, parallel to the long axis of the sleeve, thereby to pull the stent into and along the lumen of the sleeve.

In another aspect of the invention, the longitudinal forces applied to the stent as the stent advances through the constrictor and into and through the sleeve are monitored continuously by a sensor such that advancement of the stent can be terminated immediately should the force reach or exceed a predetermined level. In still another aspect of the invention, the compressed stent is loaded into one end of the sleeve and is drawn substantially along and through the full length of the sleeve to a position at a predetermined distance from the distal end of the sleeve.

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According to another aspect of the present invention there is provided a method of installing a stent in a sleeve, comprising the steps of

- i) coupling a leading end of the stent, in an expanded configuration of the stent, to an adaptor
 - ii) drawing the adaptor lengthwise within the sleeve, to pull the stent into the sleeve, and
 - iii) de-coupling the adaptor from the stent, once the stent is inside the sleeve.

Of note is the concept of gripping or clamping the stent receiving end of the sleeve so that, as the stent is pulled down the lumen of the sleeve, the resulting frictional resistance imposes on the sleeve a tensile axial stress rather than a compressive stress and behind the stent rather than in front of the advance of the stent. This facilitates advance of the stent along the sleeve lumen.

In one useful embodiment of the invention, the elongate puller is provided as a wire, and that wire is mounted to a post which serves as the puller holder. The post is fixedly mounted to the base, which extends under the device and provides a substrate for a carriage, which carries the gripper, gripper holder and constrictor. A servo-motor in the carriage serves as the translator, to move the carriage along the base away from the post, thereby to translate the sleeve past the stent, as the stent is held on the wire. A force sensor is interposed between the wire and the post, in order to monitor the tensile force in the wire. This force sensor inputs a data processor which is itself controlling the servo-motor, so that translation can be stopped if the tensile force in the wire rises too high. Conveniently, one or more sensors can be located adjacent the sleeve, at the far end, to monitor the arrival of the stent at the far end, and ensure that the stent is brought to the desired end there.

Characteristic of the invention is the adaptor which is able to engage an end of the stent when the stent is expanded, and also adapted to be released from the stent, when the stent is compressed.

In a preferred embodiment of the adaptor, a circle of hooks is provided, each of these hooks being on a long stem running hack to the elongate puller. Each of the hooks can be provided with a bight which has a radially outward-facing opening, so that the circle of hooks can be introduced inside the rim of the leading end of the stent, following which, the outward opening bight of each hook is passed through respective apertures in the cylindrical envelope of the stent, so that the point of each hook lies outside the cylindrical envelope of the stent. The circle of hooks is arranged at evenly-spaced intervals around the circumference of the leading end of the stent. In a typical laser cut cylindrical shape memory alloy stent, in its expanded configuration, there are rhombic apertures in the cylindrical envelope of the stent, and thus a circle of rhombi at the leading end of the stent. Conveniently, there is a hook corresponding to each of the rhombi in the circle of the leading end of the stent. However, one might also provide half as many hooks as there are rhombi, or some other arrangement, so long as the circle of hooks is sufficiently numerous to impose an even pull around the circumference of the leading end of the stent.

In another embodiment, the hooks can be provided as a ball on the end of each stem. With the stem in tension, and the ball radially outside the cylindrical stent envelope, the rhombus vertex catches the ball and prevents the ball passing radially inwardly through the stent envelope, until the tension on the wire is released.

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Once the stent has been pulled into the sleeve, the adaptor has to be detached from the leading end of the stent. Conveniently, the structure of the adaptor has enough column strength to allow it to be pushed towards the stent, so that the circle of hooks is itself urged towards the trailing end of the stent, thereby freeing the bight of each hook from the respective vertex of the leading end rhombus with which it is engaged. With the bight of the hook now spaced away from the rhombus vertex at the leading end of the stent, and somewhere in the middle of the rhombic space through which it passes through the cylindrical envelope of the stent, a collar surrounding the circle of stems of the hooks can be urged towards the trailing end of the stent. With the diameter of the collar sufficiently small, the stems are urged radially inwardly, themselves bringing the circle of hooks to a circular envelope with a diameter smaller than the inside diameter of the compressed stent wall. In this configuration, with the hooks no longer penetrating into the cylindrical envelope of the stent, but instead lying entirely radially within it, the way is clear for the circle of hooks to be withdrawn, axially away from the trailing end of the stent, past the leading end of the stent, and out of the sleeve. Preferably, the collar has a leading end rim which passes into the open bight of each of the hooks, and fills the bight, thereby denying any other structure the possibility of falling into the bight of each hook, as the circle of hooks is withdrawn past the leading end of the stent.

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For example, each stem could end in a loop instead of a hook, and a purse string could be threaded through apertures of the stent as well as each of said loops.

Release of such an adaptor is effected by pulling out the purse string. In such an embodiment, the stems need not be resilient and no surrounding collar is required.

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In another embodiment, a plurality of pull strings or wires are used each being threaded through two apices of a stent matrix at the leading end of the stent, these two apices being at opposite ends of a diameter across the stent lumen. Two, three,

four or more of such strings can be arranged across different diameters. On the axis, where all strings cross, a single retraction string can be looped around all the pull strings, and can lead out through the stent past the opposite, trailing end of the stent, to pull out the pull strings when their work is done.

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Alternatively, each pull string could have its own release string. Each pull string could pass through only one stent apex, or more than two apices.

An adaptor of another form entirely could be used. For example, one resembling a bottle brush might be feasible. In such an adaptor a structure along the axis of the device has a plurality of push strands which are arranged around the circumference of the axis and are resistant to longitudinal compressive stress. One end of each strand is fixed to the axial structure and the other is cantilevered radially outside it. With the fixed ends closer to the trailing end of the stent, pulling on the axial structure from in front of the leading end of the stent brings the cantilevered ends of the strands into contact with push surfaces of the stent, such as apices of its mesh. Continued pulling advances the stent by virtue of the push given to its push surfaces by the strands, now in axial compression. When it is time to release the adaptor from the stent, it can simply be urged rearwardly, past the trailing end of the stent, and the strands slip past the luminal-facing surfaces of the stent.

In a variant, the push strands could be arranged as the ribs of an umbrella with its apex remote from the leading end of the stent and its ribs engaging the stent near its leading end.

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For release of such a device, one could either pull on a release string from behind the trailing end of the stent, or push on a push shaft from in front of the leading end of the stent. Another possibility would be to use a ring which embraces the push strands, sliding the ring towards the cantilevered ends to draw them radially inwardly out of engagement with the stent.

An adapter alternately may have the general form of a chuck, which grips the leading end of the stent between two co-axial annular members, which move axially towards each other to grip the stent and away from each other to release the stent.

It will be appreciated that in the presently preferred embodiment of loading device the location of the stent relative to the base remains constant, and the translator moves the sleeve relative to the base, until the stent is in the desired location within the sleeve. However, in alternative embodiments, it could be arranged that the sleeve does not move relative to the base, but the stent is translated, relative to the base as well as the sleeve.

The invention is particularly well suited to the installation of stents which are otherwise not well adapted to installation under water in a sleeve. Thus, it is well adapted to installing encapsulated or covered stents, but can also be used to install bare stents.

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The invention is particularly attractive in respect of stents which are themselves sleeved or encapsulated. With the stent being pulled from its leading end, there is friction acting between the stent and the surfaces which are compressing it, leading to a tensile stress within the length of the stent. Some stents are not well adapted to withstand such longitudinal tensile stresses, but sleeved or encapsulated stents are often better able to resist such tensile stresses, because the encapsulation or sleeve can share the longitudinal tensile stresses imposed on the stent.

In one combination, the stent is one which carries a sleeve of expanded PTFE

(ePTFE), which is fixed to the metallic cylinder of the stent at zones corresponding to the leading and trailing ends of the stent, but not over the middle part of the length of the stent. One way of achieving this result is to place bands of ePTFE inside the cylindrical envelope of the stent, one at each end of the stent, on a mandrel which then receives the stent and the abluminal ePTFE sleeve. The assembly is sintered, causing the sleeve and bands to bond together, through the apertures of the stent cylinder, at each end of the stent. The ePTFE sleeve is relatively inextensible, and therefore supports the metal stent cylinder against elastic and plastic deformation

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which might otherwise occur under the tensile stresses imposed on the stent as it is drawn into the sleeve.

As to the loading of the stent into a sleeve, the sleeve is itself elastically deformable, and, in one loading technique, will be subject to tensile stresses during the stent installation process. These forces will produce tensile strain in the sleeve. Unless allowance is made for this strain in the installation operation, then the actual position of the stent in the sleeve will not correspond to the desired position. To compensate for the effect of strain in the sleeve, the speed of translation is reduced as the stent approaches the desired end point of its travel within the sleeve. A laser beam extending across the path of the sleeve can serve to detect the presence or absence of the sleeve, thereby detecting whether the sleeve is still strained and, in effect, establishing whether the actual position of the far end surface of the sleeve corresponds to the desired location. A magnetic inductance sensor detects the arrival of the stent being translated with the sleeve at the desired location near the distal end of the sleeve. Data from these two sensors, delivered to the data processor, allow the data processor to terminate the pulling process at a point which places the stent at a desired distance from the far end surface of the sleeve.

For a better understanding of the present invention, and to show more clearly how the same may be carried into effect, reference will now be made, by way of example, to the accompanying drawings.

Brief Description of the Drawings

Figure 1 is a longitudinal diametrical section through the stent, adaptor, gripper and constrictor of the device of the present invention;

Figure 2 is a longitudinal diametrical section through the adaptor of the device, showing its internal structure;

Figure 3 shows a sleeved stent with a leading end engaged by the adaptor;

Figure 4 is a longitudinal section through a first embodiment of the device of the present invention, showing it in an initial configuration; and

Figure 5 is a section corresponding to that of Figure 4, showing the device in a final configuration.

Figure 6 is a longitudinal diametrical section of a second embodiment of stent loading machine engaging a stent which has a plurality of beads arranged around its leading end;

Figure 7 is a section as in Figure 6, showing a later stage in the loading of the stent;

Figure 8 is a diagrammatic illustration of a stent threaded with wires to facilitate drawing of the stent through a constricting device to compress the stent and lead it into an end of the sleeve;

Figure 9 is a highly diagrammatic illustration of the end of the stent as shown in Figure 8, as seen from the right end of Figure 8;

Figure 10 is a diagrammatic illustration of the apparatus for supporting the sleeve as the stent is drawn through the constrictor and into the sleeve; and

Figure 11 is an illustration of the enlarged open end of the constrictor showing a stent partially drawn into the constrictor, the stent having been reinforced with an internal support tube.

Detailed Description

Referring first to Figure 1, there is shown a stent 10 in its expanded configuration, and with a leading end 12 and a trailing end 14, an abluminal surface 16 and a luminal surface 18. A circle of hooks 20 engages the leading end 12 of the stent. Each of the hooks 20 has a long stem 22, and all of these stems extend to a root portion 24 located close to a long axis 26 of the device in general. Over the circle of root portions 24 of the stems 22 is a collar 28 which is slidable over the stems 22, up to the hooks 20.

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In other embodiments, detent surfaces other than hooks could be used. For example, a little ball of the end of each stem 22 might be effective, depending on the design of the locations in the stent where the stems are to be coupled.

An elongate puller in the form of a wire 30 is attached to the portions 24 of the stems 22 and extends through and beyond the collar 28. The structure of hooks 20, stems 22, and collar 28 constitute an adaptor 41 which couples the trailing end 43 of the pulling wire 30 to the leading end of the stent 10.

The sleeve 32 into which the stent is to be loaded is supported to receive the pulling wire 30 that extends through the lumen 33 of the sleeve. In the embodiment shown in FIGS. 1-5, a receiving end 50 of the sleeve 32 is gripped in a gripper chuck 34. The chuck 34 is in fixed position or relationship to a constrictor 36 which is in the form of a block 38 of large thermal capacity, which itself defines a tapered, conveniently conical, constricting surface 40 with a wide end 42 and a narrow end 44. At this narrow end 44 is a shoulder surface 46 at the end of a bore surface 48 which snugly receives the receiving end 50 of the sleeve 32. The sleeve has a long axis corresponding to the axis 26 of the device in general. As will be described, the stent will be compressed and drawn into and through the sleeve 32 until the stent is located in a predetermined position at the opposite end of the sleeve. The receiving end 50 of the sleeve thus will serve as the proximal end of the delivery device and the opposite end of the sleeve will serve as the distal end to be inserted into the patient. When the stent is positioned in the sleeve in readiness for deployment in the patient, it will be located near the distal end of the sleeve.

Also to be seen in Figure 1 is a mandrel 52 which has a tapered conical surface 54 which complements the tapered conical surface 40 of the constrictor block 38, so that advancement of the mandrel 52 along the axis 26, into the conical cavity of the constrictor block 38 leads to the definition of a conical annular space through which the hook stems 22 extend.

Conceivably, the constrictor could oscillate axially to assist advance of the stent down to its narrow end. If so, it may be helpful to provide the taper surfaces with a "one-way" structure of steps and terraces, like the underside of a cross-country ski, so that the constrictor can slip backwards relatively easily but on its forward stroke towards the leading end of the stent, tends to carry the stent with it. Alternatively or additionally the cone could be coated with a slippery material to assist stent advance.

Not shown in Figure 1, for reasons of clarity, are the details of the surrounding structure of the device. (For this, see Figures 4 and 5 and the description below, in relation to those figures).

In operation, a carriage which carries the gripper chuck 34 and constrictor 36 and mandrel 52 is caused to move to the right as shown in Figure 1, while the pulling wire 30 is held fast. As will be appreciated, this has the effect of drawing the hooks 20 down the annular space adjacent the tapered conical surface 40 of the block 38, and the hooks carry with them the leading end 12 of the stent 10. Thus, progressively, the full length of the stent 10 is drawn down the cone and into the receiving end 50 of the sleeve 32. Further progress of the carriage to the right in Figure 1 pulls the stent 10 along the full length of the sleeve 32.

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Turning now to Figure 2, the internal construction of the adaptor is visible. A welding bead of fillet 64 fixes the root zones 24 of the stems 22 to the pulling wire 30. The stems 22 are surrounded by a moving collar 62. The fixed sleeve 60 is attached by a welding bead 64 to the pulling wire 30. Attached to the moving collar 62, by a further welding bead 66, is a collar ring 68 having a leading end rim '70. The outside diameter of the collar ring 68 is somewhat smaller than the outside diameter of the moving collar 62. Inside the moving collar 62 is a plurality of apertured rings 72. In the present embodiment, there are twelve hooks 20 and each of the rings 72 has a circle of 12 apertures, each to receive a different one of the stems 22. These 20 apertured rings '72 serve to maintain the spaced even circular arrangement of the long stems 22 and help the stems resist buckling in response to a compressive load.

In use, the moving collar 62 is brought to a proximal location as shown in Figure 2. The stems 22 are provided with a relaxed disposition in which they curve radially outwardly, so that the hooks 20 are on a circular envelope of relatively large diameter. Sliding the collar 62 to the right hand side in Figure 2 brings the rim 70 of the collar ring 68 into the bight 74 of each of the hooks 20. The stems 72 deform elastically, and the circle on which the hooks 20 is located is now a circle of radius corresponding to the radius of the collar ring 68. This is small enough, relative to the inside diameter of the stent cylinder, to allow the hooks 20 to pass along the cylindrical cavity within the stent, without engaging on any part of the apertured wall of the stent, and the presence of the collar ring 68 inside the bight 74 of each hook itself prevents any other structure from passing into the bight of the hook.

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Figure 3 shows the circle of hooks 20 each engaged with a different rhombic aperture 80 at the leading end of the stent 10. In fact, each rhombic aperture has a vertex 82 closest to, the leading end 12 of the stent 10 and it is with this vertex 82 that each hook 20 is engaged, for pulling the stent 10 into the sleeve 32.

For coupling the hooks 20 to the stent 10, the collar 62 is slid tip to the hooks 20 and then the circle of hooks 20 is introduced into the leading end of the stent, to bring each of the hooks 20 into a position more or less central within its corresponding rhombic aperture 80. Then the collar 62 is slid proximally back towards the Figure 2 position, allowing the hooks 20 to expand radially outwardly by resilient elastic deformation of the stems 22, until each hook 20 penetrates through the cylindrical envelope of the stent 10 at one of the rhombic apertures 80, with each of the twelve hooks, 20 extending through a different one of the circle of twelve rhombic apertures 80 at the leading end of the stent 10.

Reverting again to Figure 1, the sleeve 32 destined to receive the stent 10 is gripped in chuck 34 and the wire 30 is, then passed through the constrictor block 38 and sleeve 32, until its leading end 39 can be made fast to a puller holder, ready for pulling the stent 10 into the sleeve 32.

Now looking at Figure 4, one sees the leading end 39 of the pulling wire 30 made fast to a post 90 on a base 92 and with a force sensor 94 in the post 90 and arranged to monitor tensile force in the pulling wire 30. The sensor 94 inputs, along lead 96, a data processor 98.

On the base 92 is a carriage 100 mounted for linear translational movement towards and away from the post 90. Within the carriage 100 is a servo motor 102 controlled, via lead 104, by the data processor 98. The carriage 100 carries the constrictor 36, mandrel 52 and sleeve gripper 34. Further, the sleeve 32 is supported from below on a beam 106 which extends from the carriage 100 the full length of the sleeve 32, up to a peninsular block 108 supported above the base 92 by an undercarriage 110. The

peninsular block 108 travels with the carriage 100, when the servo motor 102 is actuated.

Mounted on the peninsular block 108, which supports the far end of the sleeve 32, is a plurality of sensor elements, in particular, an optical sensor 112 and a magnetic induction sensor 114. The inductive sensor 114 detects the arrival, at the distal 37 of the sleeve 32, of the stent 10, and to signal that arrival to the data processor 98, through lead 118. The optical sensor 112 passes a laser beam across the locus of the sleeve 32 to sense the end of the sleeve and inform the data processor through lead 116, so that the stent 10 can be brought to a precisely specified location in the sleeve 32 relative to the sleeve end. The sleeve 32 is fixed against movement relative to the peninsular block 108, but only at its receiving (proximal) end 50, so that tensile stresses in the sleeve 32 can alter the position of its distal 37 relative to the sensor 112. Provision of sensor 112 therefore can inform the data processor when the tensile strain in the sleeve 32 has fallen below a specified low value, corresponding to arrival of the stent at the desired end position, and enabling the pulling process to be terminated to lace the stent at just the correct desired distance from the distal end of the sleeve. Readers will appreciate that elastic deformation of the synthetic polymeric materials of stent sleeves is a time-dependent phenomenon, so time has to be allowed, at the end of the pulling process, for the sleeve to relax into its unstressed length.

Not shown in Figure 4 are the arrangements for temperature control of the block 38 of the constrictor 36. Those skilled in the art will not find it difficult to install arrangements to bring the block 38 to the temperature prescribed for optimum installation of the stent.

Figure 4 shows the device before stent installation begins, and Figure 5 shows the device once stent installation is complete. With reference to Figures 4 and 5, the stent installation process proceeds as follows.

With the pulling wire coupled to the stent, and installed as described above with reference to Figure 1, the carriage 100 is moved to the right in Figure 4, so as to bring

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the leading end 12 of the stent 10 to the wide end 42 of the conical constricting surface 40. The mandrel 52 is advanced to the left in Figure 4, along a key way 120 until it fits within the block 38 to complement the block conical surface 40. Further translation of the carriage 100, in the direction of arrow F, has the effect of carrying the constrictor block 36 along the length of the stent 10, from the leading end 12 towards the trailing end 14, so that the length of the stent is drawn down the conical surface 40 and into the receiving end 50 of the sleeve 32. Should at any time the tensile stress in the pulling wire 30 exceed a pre-determined limit, as measured by force detector 94, then the data processor 98 arrests movement of the carriage by terminating the drive power to the servo motor 102.

As drawing of the stent 10 along the sleeve 32 proceeds, so the receiving end of the sleeve experiences a tensile stress. Note, however, that the sleeve ahead of the stent is not under any axial stress so that stresses in the sleeve at the leading end of the stent are at a minimum, facilitating progress of the stent along the lumen of the sleeve. Effectively, with the sleeve in tension, one is pulling a sleeve over the stent, as opposed to pushing a sleeve over the stent. Towards the end of the pulling process, as shown in Figure 5, substantially the full length of the sleeve 32 is under tensile stress, leading to an increase in its length. However, as explained above, the information which the data processor receives from sensors 112, 114 allows it to translate the sleeve 32 to a position relative to the stent 10 which sets the stent 10 at exactly the desired distance from the end surface of the sleeve 32. At that point, further translation of the carriage 100 is stopped, and the installation process is complete. At: this point, the device can be switched off, and the sleeve 32 removed from it, with the stent 10 in place, yet still connected by the adaptor hook ring to the pulling wire 30.

Reverting now to what is shown in Figures 2 and 3, it will be appreciated that, to remove the circle of hooks 20 from the installed stent 10, the tension has to be taken off the pulling wire 30, and the hooks 20 allowed to relax into the center of the rhombic apertures of the stent 10 in which they are engaged. Then, a movement of the collar 62, up to the hooks 20, has the effect of bringing the rim 70 of the collar ring 68 into the bight 74 of each of the hooks 20, and bringing the diameter of the circular

hooks 20 down to a diameter small enough to allow all of the hooks to be drawn out from the cylindrical cavity of the stent, without fouling the cylindrical envelope of the stent.

At this point, the stent and sleeve stand alone, and can be passed forward for the further steps of the process of constructing the stent delivery system.

Important to note is that the distal end of the sleeve is not engaged mechanically by any component of the loading system. The end can therefore be polished in an earlier process step, and does not need to be cut or polished after loading. This is important, because the steps of cutting and polishing could conceivably give rise to the undesirable presence of loose particles inside the sleeve at its distal end. The other end of the sleeve, which has been held in the gripper chuck 34, is the proximal end of the sleeve in use. This can be cut, if necessary, after the stent has been loaded.

Figure. 6 shows a leading end portion of another embodiment of stent 112. At the end vertex of each cell in the first circumferential ring of cells of the stent, the material of the stent matrix is continued into an extending portion 174 with a width comparable to its thickness dimension so that, in cross-section, it is more or less square. On each such square section spigot 174 is mounted a spherical Nitinol bead 176 which has a through bore on a diameter of the bead, to receive the spigot 174. The Nitinol bead 176 is welded to the spigot 174. It will be appreciated that, by virtue of the rounded surface and greater thickness of the sphere 176 relative to the struts 154, the free vertices defining the end of the stent, and the end of each cell in the end ring of cells of the stent, is less likely to cause trauma in the bodily tissue in which the end vertices is embedded, than if the spheres 176 were absent.

Furthermore, as shown in Fig. 6, the ring of beads 176 brings advantages when it comes to loading the stent onto a delivery system, and keeping control of the stent while the stent is being deployed into the body from the delivery system. Specifically, the ring of relatively thick beads 176 provides a point of purchase for gripping surfaces to impose forces on the stent, while it is being loaded into a delivery system,

and while it is being deployed from that delivery system. In one example, the beads 176 could be gripped between circumferential surfaces, one inside the stent annulus and one outside the stent annulus, with a spacing between such co-axial surfaces which is wide enough to receive the thickness of the stent matrix, but does grip the spheres 176 on each side of the thickness of the stent matrix.

Viewing Figures 6 and 7, the stent 112 is shown schematically within the truncated cone of a loading mandrel 190, with its .1eading end at the narrow end of the cone, tipped by the beads 176. Within the leading end of the stent is a loading rod 192 with a somewhat larger diameter head 194, the transition from the head 194 to the cylindrical portion 196 of the rod 192 being accomplished by an arcuate shoulder surface 198. The concave outer surface of the shoulder 198 has a curvature which corresponds to the curvature of the beads 176.

Beyond the narrow end of the truncated cone 190 is a gripping sleeve 200 which has at its gripping end 202 an arcuate gripping shoulder 204, also having a curvature corresponding to that of the spherical surface of the bead 176.

As can be seen from Figure 7, drawing the gripping rod 192 down on to the beads 176 achieves an entrapment of the beads 176 in an annulus defined by the gripping shoulders 198 and 204. With the position of the gripping rod 192 maintained close to the gripping end 202 of the gripping sleeve 200, further pulling down of the gripping rod 192, away from the truncated cone 190, permits the advancement of the stent 112 into the cylindrical space shown in Figure 7, within the block 206.

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The block 206 receives a sleeve 208 in which the stent 112 is to be housed, in a delivery system for placing the stent 112 at a desired location within the body, for location, a catheter. Continued downward pulling on the gripper rod 192, beyond the position shown in Figure 7, can carry the stent 112 fully inside the sleeve 208 of the catheter delivery system. Once the stent 112 is within the sleeve, the gripping sleeve 200 can be withdrawn forwardly, i.e. downwardly in the Figure 7 view, while the gripper rod 192 can be withdrawn rearwardly from the stent, i.e. upwardly as shown in Figure 7 and back past the trailing end of the stent. Alternatively, once the gripping

sleeve 200 is withdrawn, it may be possible to withdraw the gripper rod 192 forwardly, given a degree of resilience in the sleeve 208 to allow the enlarged head 194 to slide past the beads 176 at the leading end of the stent.

Figures 8-11 illustrate, diagrammatically, another technique and apparatus for 5 compressing the stent and loading it into the distal end of the sleeve of the delivery device. In this technique, one or more wires, such as slender, superelastic nitinol wires, may be attached to the rhombi or diamonds, at the proximal end of the stent. Figure 8 shows two such wires 310, 312. The wire 310 is passed through a pair of diametrically opposite diamonds 308a, 308b and another wire 312 may be passed through another pair of diametrically opposite diamonds 308c, 308d. The portions of the wires 310, 312 that span the cross-section of the stent intersect at a point P located at or near the longitudinal axis of the stent. The free tails of the wires 310, 312 may be wrapped together, as with tape, and threaded through a funnel-like constrictor 314. The constrictor has an enlarged inlet end 316 and a narrowed outlet 15 end 318 that opens into a fitting 320 adapted to engage the distal end 322 of the sleeve 324 into which the stent is to be loaded. The tails of the wires 310, 312 are passed through the sleeve 324 and exit out of the proximal end 326 of the sleeve. The device includes a clamp 328 that is separable to receive the sleeve 324. The clamp may be of a two-part construction having thumb screws 330 to enable the clamp parts to be secured together to clamp the sleeve 324. The clamp 328 is in a fixed position with respect to the constrictor 314, as by securing them to a common frame (not shown). The device also includes a trough-like support 332 that extends between the clamp 328 and the fitting 320 to support the portion of the sleeve 324 that extends between those components. With the system set-up as illustrated in FIG. 10, the stent can be drawn through the constrictor 314 and into the lumen of the sleeve 324 simply by pulling on the tails of the wires 310, 312. In this embodiment, the sleeve 34 may be transparent or translucent sufficiently to enable the stent to be visually observed as it advances into the distal end of the sleeve 324. Under such visual observation, the stent may be drawn into the distal end of the sleeve 324 to the 30 precise location desired. The pull wires 310, 312 then may be withdrawn by pulling on one of the tails of each of the wires.

In a modification of the foregoing, an additional retraction wire 334 can be employed to enable withdrawal of the pull wires out of the distal end of the sleeve 324. In that embodiment, a retraction wire 334, constructed similarly to the pull wires 310, 312 is inserted through the distal end of the stent, out through the proximal end, and is looped about the crossed region P of the pull wires 310, 312, the end of the retraction wire 34 being passed back out through the distal end of the stent. In this embodiment, after the stent has been positioned at the intended location within the sleeve 324, the pull wires 310, 312 are withdrawn by pulling on the retraction wire in a distal direction. The retraction wire 334, being looped about the crossed pull wires, draws the pull wires, at their bight, through the sleeve 324, the stent, and the constrictor 314.

It may be desirable to provide internal support for the stent, as the stent is drawn through the constrictor. To that end, a polyimide support tube 336 can be inserted into the distal end of the stent as the stent is advanced into the constrictor. The polyimide tube is sufficiently stiff to provide luminal support for the stent as the stent is constricted and compressed to its smaller diameter. Additionally, after the stent has been loaded into the distal end of the sleeve 324, the presence of the inner support tube 336 assures that a guidewire can be passed through the sleeve 324 and the stent in order to facilitate navigation of the delivery device through the patient's vasculature. It may be noted that the polyimide tube 336, although providing internal support for the stent, nevertheless enables the retraction wire 334 as well as the pull wires 310, 312 to slide between the inner lumen of the stent and the support tube 336 to permit their removal.

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Industrial Applicability

The invention contributes to the art of stents by providing improvements in methods of, and apparatus for, loading a stent into a sleeve of a delivery system in a way which can be controlled by a data processor. Such automation furthers the objectives of consistency and quality control. Data for particular designs of stent and sleeve can be stored in the data processor, with the corresponding parameters, especially the limiting load on the pulling system, in order that the data processor shall stop pulling when the limit load is exceeded. Thus, when a batch of stents is to be loaded, the

data processor will drive the system in accordance with a constant program using the stored data.

The system described above has been found to be able to handle consistently and reliably stents of a relatively large diameter too large for manual loading. Sleeving the stent in a gaseous environment furthers the object of sterility. The invention allows ready adaptation, to enable sleeving at any desired non-ambient temperature.

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CLAIMS

- 1. A device for compressing and pulling into the sleeve (32) of a stent delivery system, in a pull direction along a long axis, a self-expanding stent (10) which has a long axis which is co-axial with that of the delivery system, the pulling device including
 - i. a funnel to lead the stent into the sleeve, the funnel having a relatively large receiving end and a relatively small delivery end, said delivery end including an annular shoulder against which abuts, in use of the device, a receiving end of said sleeve and
 - ii. pull structure (30) to be received within the sleeve, said pull structure defining a plurality of pull points (20), arranged around an annulus, for engaging the stent in an annular locus, around its long axis; the device being characterized by:
 - iii. a clamp to restrain the sleeve from following the stent, as the stent is pulled into the receiving end of the sleeve; and
 - iv. means (62) to release the pull structure from the stent, after the full length of the stent from a leading end thereof through to a trailing end thereof has been pulled down the funnel and into the sleeve, by a movement of the pull structure from a pulling disposition to a release disposition in which the pull structure is no longer in abutment with the stent, thus permitting the pull structure to be withdrawn from within the sleeve, while the stent remains within the sleeve.
- 2. Device as claimed in claim 1 wherein the pull structure comprises a plurality of strands (22) each having a length direction component along said long axis, and the plurality being distributed around the axis.

- 3. Device as claimed in claim 2 wherein the strands are pull strings (22) thrown into tension when pulling the stent into the sleeve.
- 4. Device as claimed in claim 3 wherein each pull string is a loop which extends from a proximal end outside the sleeve to a distal end corresponding to a pull point and then returns to said proximal end.
 - 5. Device as claimed in claim 3 or 4 wherein each pull string has a detent surface (20) to abut the stent.
 - 6. Device as claimed in claim 5 wherein the detent surface is defined by a hook (20).
- 7. Device as claimed in claim 2 wherein the strands are push elements, thrown into compression when pulling the stent into the sleeve, each such strand being attached at a proximal end to a pull shaft and extending from said proximal end in said pull direction to a free distal end which, during pulling, abuts a push surface of the stent which faces away from the pull direction.
- 20 8. Device as claimed in claim 7 wherein the pull shaft and push elements together have the overall form of a bottle brush.
 - 9. Device as claimed in claim 7 wherein the pull shaft and push elements together have the overall form of an umbrella shaft and ribs.
 - Device as claimed in any one of the preceding claims, in which the release means is actuated from a point beyond the stent, and withdraws the pulling structure past the trailing end of the stent.
- 30 11. Device as claimed in claim 10, the pulling structure comprising a plurality of pulling string loops, and the release means comprising at least one release string.

- 12. Device as claimed in claim 11 wherein each pulling string has its own release string.
- 13. Device as claimed in claim 11 wherein a single release string is coupled to each of the pulling strings, to pull all of the pulling strings away from the stent simultaneously.
- 14. Device as claimed in claims 7, 8 or 9 wherein the release structure uses the pull shaft (30) to push the push element in a direction opposite to the pulling direction, to draw the distal ends of the push elements away from abutment with the respective push surfaces of the stent, so that the luminal surface of the compressed stent may urge the distal ends of the push elements radially inwardly, into the stent lumen space.
- 15. Device as claimed in claim 7, 8 or 9 wherein the release structure comprises a release string for pulling the pull shaft in a direction opposite to the stent loading pull direction, to pull the pull structure out of the stent lumen past the trailing end of the stent.
- Device as claimed in any one of the preceding claims, wherein the release means comprises a ring (62) to urge the pull points radially inwardly, for bringing them into the luminal space of the compressed stent.
- 17. Device as claimed in claim 16 wherein the pull structure comprises a plurality of resilient strands (22) arranged each with a component parallel to each other and spaced around the axis, and each of which has a proximal end at the axis of the structure and a distal end at which is one of said pull points (20), and has a relaxed disposition in which the locus of said pull points defines a circle with a radius larger than the circle defined by the respective proximal ends, the ring release means (62) surrounding the resilient strands and acting, as the ring moves along the lengths of the resilient strands from their proximal ends towards their distal ends to urge the distal end pull points radially inwardly.

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- 18. Device as claimed in claim 16 wherein the release means ring is a string which functions as a draw string or purse string.
- 19. Device as claimed in any one of the preceding claims wherein the pull structure comprises a first annular element and the release means comprises a second annular element, the first and second annular elements comprising a chuck.
 - 20. Device as claimed in claim 19 wherein the first annular element lies radially inside the second annular element and so abuts the stent from its luminal surface.
 - 21. Device as claimed in claim 20 wherein the first annular element defines a frusto-conical surface which tapers radially inwardly in the pulling direction, the pull points being found within said frusto-conical surface.
- .22. Device as claimed in claim 19, 20 or 21, in which the first and second annular elements move relative to each other, along the axis, between an approximated disposition for gripping the stent therebetween and a less approximated position for releasing the stent from the chuck.
 - 23. Device as claimed in any one of the preceding claims, including a puller (102), to impose a load on the sleeve and the pull structure to advance the stent into the sleeve.
- 25 24. Device as claimed in claim 23 including a load sensor (94) and a device to protect against the imposition on the stent by the puller a stress during loading which exceeds a limit stress.
- Device as claimed in claim 23 or 24, including an end point sensor (114) which can detect the arrival of the stent at a desired location in the sleeve and then output a stop signal to the puller.

- 26). Device as claimed in any one of the preceding claims, arranged to pull a stent into a sleeve in a direction such that the leading end of the stent is to become the distal end of the stent in surgery.
- 5 27. Device as claimed in any one of the preceding claims, and including a funnel block (36) to guide the stent and compress the stent prior to its advancement in the receiving end of the sleeve.
- Device as claimed in any one of the preceding claims and including a mandrel
 (52) to fit inside the funnel, to resist any unwanted radially-inward buckling of
 the stent between its leading and trailing ends, as the stent advances along the
 funnel towards the narrow end of the funnel.
- 29. Device as claimed in claim 27 or 28 and including means to establish in the funnel a temperature other than ambient temperature.
 - 30. Device as claimed in any one of claims 27 to 29 including means to oscillate the portion in directions parallel to said long axis, to assist progress of the stent towards the narrow end of the funnel.
 - 31. Device as claimed in any one of claims 27 to 30 in which the surface of the funnel over which the stent slides is coated to assist such sliding.
- 32. Device as claimed in claim 31, in which said funnel surface is presented as a multiplicity of relatively wide annular terraces separated by annular lines from relatively narrow annular steps, and so arranged that each annular line has a said step immediately to its conically narrow side and a said terrace immediately to its conically more open side, thereby to resist movement of the stent towards the open end of the funnel portion.
 - 33. A device for installing a stent (10) in a sleeve (32):

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the stent having luminal (18) and abluminal surfaces (16), a leading end (12) and a trailing end (14) and having a capacity to expand radially from a compressed configuration to an expanded configuration;

the sleeve (32) having a long axis (26), a lumen (33), a receiving end (35) and a far end (37); the device comprising:

an elongate puller (30), to be received in the sleeve and having a leading end (39) and a trailing end (43)

an adaptor (41), for coupling the trailing end of the puller to the leading end of the stent when the stent is in it expanded configuration and for decoupling the puller from the stent when the stent is in its compressed configuration;

a gripper (34), to grip the receiving end of the sleeve;

a constrictor (40), to guide the leading end of the stent into the receiving end of the sleeve;

a base (92);

a puller holder (90), to transfer stresses between the puller and the base;

a gripper holder (100), to transfer stresses between the gripper and the base; and

a translator (102), to increase the distance between the puller and the gripper, parallel to the long axis of the sleeve, thereby to pull the stent into and along the lumen of the sleeve.

- 34. Device as claimed in claim 33, wherein the puller is a wire (30).
- 35. Device as claimed in claim 33 or 34, wherein the adaptor has a long axis and comprises a plurality of hooks (20) each on its own stem (22), the stem being arranged in a circle surrounding its long axis and extending along said axis.
- 36. Device as claimed in claim 35, wherein each of the hooks has a radially outwardly-open bight (74).

- 37. Device as claimed in claim 35 or 36, wherein the adaptor includes a collar (62) which is slidable on said axis, along the stems and up to the hooks, to draw the hooks towards said axis.
- Device as claimed in claim 37, as dependent on claim 36, wherein the collar has a leading end (68) with a leading edge (70) which is received in the bight of each of the open hooks to close the bight.
- Device as claimed in any one of the preceding claims, wherein the adaptor is fabricated from a shape memory alloy.
 - 40. Device as claimed in any one of the preceding claims, wherein the gripper is a chuck (34).
- Device as claimed in any one of claims 33 to 40, wherein the gripper and the constrictor are mounted to move together.
 - 42. Device as claimed in any one of preceding claims 33 to 41, wherein the gripper holder comprises a carriage (100), which the translator moves relative to the base.
 - 43. Device as claimed in claim 42 which includes a load sensor (94) mounted in the puller holder to sense the load between the puller and the base.
- Device as claimed in claim 43 which includes a force data processor (98) which receives data from the force sensor and can modify the performance of the translator in dependence on such data, in accordance with a force program.
- Device as claimed in any one of the preceding claims, wherein the constrictor comprises a tapered surface (40) to compress the abluminal surface of the stent.

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- 46. Device as claimed in any one of claims 33 to 45, wherein the constrictor includes a mandrel (52) to support the luminal surface of the stent.
- 47. Device as claimed in any one of claims 33 to 46, wherein the constrictor includes structure (38) to bring the temperature of the stent to a desired non-ambient temperature.
 - 48. Device as claimed in any one of claims 33 to 47, including a stent sensor (114) at a zone within the length of the sleeve where the stent is desired to be, said stent sensor inputting the translation data processor, to slow the movement of the translator.
 - 49. Device as claimed in any one of the preceding claims, including a sleeve far end presence sensor, to monitor tensile strain in the sleeve, and arrest translation of the sleeve when the strain is less than a predetermined low limit.
 - 50 A method of installing a stent in a sleeve, comprising the steps of
 - i) coupling a leading end of the stent, in an expanded configuration of the stent, to an adaptor
 - ii) drawing the adaptor lengthwise within the sleeve, to pull the stent into the sleeve, and
 - iii) de-coupling the adaptor from the stent, once the stent is inside the sleeve.
 - 51. Method according to claim 50, including the step of coupling the adaptor to the stent at a plurality of push points spaced around the circumference of the leading end of the stent.
- Method according to claim 51, including the step of providing the adaptor as a plurality of strands, each to engage with different ones of the push points.

- 53. Method according to claim 52, wherein each of the strands is a pull string, and each of the pull strings is associated with the push points in one-to-one correspondence.
- 5 54. Method according to claim 52, wherein each of the strands is a pull string which is looped around a pair of push points, the members of the pair being diametrically apart on the leading end of the stent.
- 55. Method according to claim 52, 53 or 54, including the step of pulling the strands off the stent from a point beyond the trailing end of the stent, to decouple the adaptor from the stent once the stent is inside the sleeve.
 - 56. Method according to claim 55, including the step of threading a release string around each of the adaptor strands, whereby pulling on the release string pulls all the strands off the leading end of the stent.
 - 57. Method according to any one of claims 50 to 54 including the step of pushing the strands off the stent from a point beyond the leading end of the stent.
- Method according to any one of claims 50 to 57 including the step of sensing a load imposed on the stent when drawing the stent into the sleeve, and arresting the drawing movement when a pre-set limit load is reached.
 - 59. Method according to any one of claims 50 to 58, including the step of defining an end position of the stent relative to the sleeve, sensing the approach of the stent to said end position, and arresting the drawing movement when the end position is sensed.
- 60. Method according to any one of claims 50 to 59, including the step of maintaining the stent at a temperature other than ambient temperature, during the drawing step.

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- 61. Method according to any one of claims 50 to 60, including the step of pulling the stent down a funnel into the sleeve.
- 62. Method according to claim 61 including the step of oscillating the funnel along its long axis, to facilitate progress of the stent towards the narrow end of the funnel.
- 63. Method according to any one of claims 50 to 62 including the step of restraining the sleeve at a stent-receiving end thereof, so that friction forces between the stent and the sleeve, as the stent is introduced into the sleeve, impose on the sleeve an axial tensile stress.
- 64. An apparatus for radially compressing a self-expanding stent and for loading the stent into a delivery sleeve comprising:

a frame;

a constrictor mounted on the frame, the constrictor having a relatively large cross-sectional inlet and a relatively small cross-sectional outlet, the region between the inlet and outlet defining a progressively smaller cross-section, the outlet being engageable with the open end of the sleeve;

a pull wire mounted at one end to the frame and at its other end detachably connectible to an end of the stent;

the pull wire and constrictor being mounted to the frame for relative movement in which the stent and constrictor may be drawn into engagement with each other;

a drive motor for effecting said relative movement;

a sensor associated with the pull wire to sense the degree of tensile force applied to the stent as a consequence of the stent's engagement with the constrictor or sleeve;

a computer having a processor programmed to control the operation of the motor to effect such relative movement, the program being responsive to a signal from the sensor of a force at or above a predetermined threshold to stop operation of the motor.

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65. An apparatus for radially compressing a self-expanding stent and for loading the stent into a delivery sleeve comprising:

a frame:

a constrictor mounted on the frame, the constrictor having an enlarged inlet receptive to an expanded stent and a smaller outlet, the constrictor being adapted to progressively compress the stent as it is drawn from the inlet to the outlet, the outlet being engageable with the open end of the sleeve;

a pull wire mounted at one end to the frame at the other end being detachably connectible to an end of the stent;

the pull wire and constrictor being mounted to the frame for relative movement in which the stent and constrictor are drawn into engagement with each other;

a support engageable with the distal end of the sleeve, the constrictor and sleeve support being configured to support the sleeve to enable the pull wire to extend into the distal end of the sleeve, through the sleeve and the constrictor;

the drive motor for effecting such relative movement;

a computer having a processor programmed to control the operation of the motor to effect said relative movement;

a sensor for sensing when the distal end of the sleeve has reached a predetermined location;

a second sensor for sensing when the compressed stent has been drawn through the sleeve to a location that is a predetermined distance from the end of the sleeve;

the computer program being responsive from the signals of said sensors to stop operation of the motor when the stent has reached said predetermined location.

66. A method for radially compressing a self-expanding stent and for loading the stent into a delivery sleeve comprising:

providing a constrictor having a relatively large cross-sectional inlet and a relatively small cross-sectional outlet, the region between the inlet and outlet defining a progressively smaller cross-section;

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supporting an open end of the sleeve in registry with the outlet;
effecting relative movement between the stent and the constrictor and
sleeve to cause advancement of the stent through the constrictor and into and
through the sleeve and, while effecting such relative movement, sensing the
force applied to the stent and terminating said relative movement when said
force exceeds a predetermined level.

- 67. In a method for drawing a stent through a constrictor to radially compress the stent and for drawing the stent from the constrictor into a sleeve, the improvement comprising monitoring the degree of axially oriented force applied to the stent during such movement and terminating the relative movement when the force exceeds a predetermined level.
- 68. A method for radially compressing a self-expanding stent and for loading the stent into a delivery sleeve comprising:

drawing the stent through a constrictor having an enlarged inlet receptive to the stent and a smaller outlet thereby to progressively compress the stent as it is drawn from the inlet to the outlet;

locating an open end of the sleeve in registry with the outlet of the constrictor;

drawing the stent from the constrictor into the sleeve and continuing to draw the stent toward the opposite end of the sleeve;

sensing the position of the opposite end of the sleeve;

sensing the position of the stent within the sleeve;

comparing the sensed positions of the end of the sleeve and the position of the stent within the sleeve;

terminating further advancement of the stent within the sleeve when the stent has reached a predetermined location with respect to the end of the sleeve.

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69. A method as defined in claim 68 wherein the end of the sleeve is sensed optically and the location of the stent within the sleeve is sensed by an electromagnetic coupling.

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70. A method for drawing a self-expanding stent through a constricting device and into a sleeve comprising:

attaching at least one wire to an end of the stent at substantially diametrically opposed locations;

orienting the open end of the sleeve with the outlet of the constrictor; inserting the pull wire through the constrictor and into and through the sleeve and thereafter pulling the pull wire to draw the stent through the constrictor so as to compress the stent to a smaller diameter;

continuing pulling the stent from the constrictor into the distal end of the sleeve;

wherein the sleeve is sufficiently transparent or translucent to enable the stent to be visibly observed through the wall of the sleeve; and

continuing to draw the stent through the sleeve until the stent has reached a predetermined position within the sleeve as observed through the sleeve.

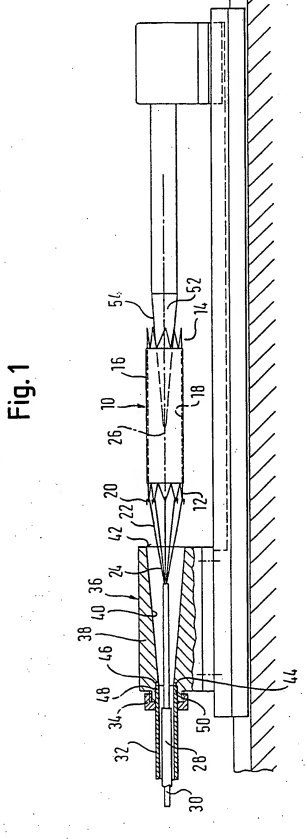
- 71. A method as defined in claim 70 wherein a plurality of said pull wires are attached to the end of the stent, the wires crossing each other at or adjacent the longitudinal axis of the stent.
- 72. A method as defined in claim 71 further comprising:

before inserting the pull wires into the constrictor, placing a retraction wire through the stent, looped about the crossing point of the pull wires and then back out the second end of the stent; and

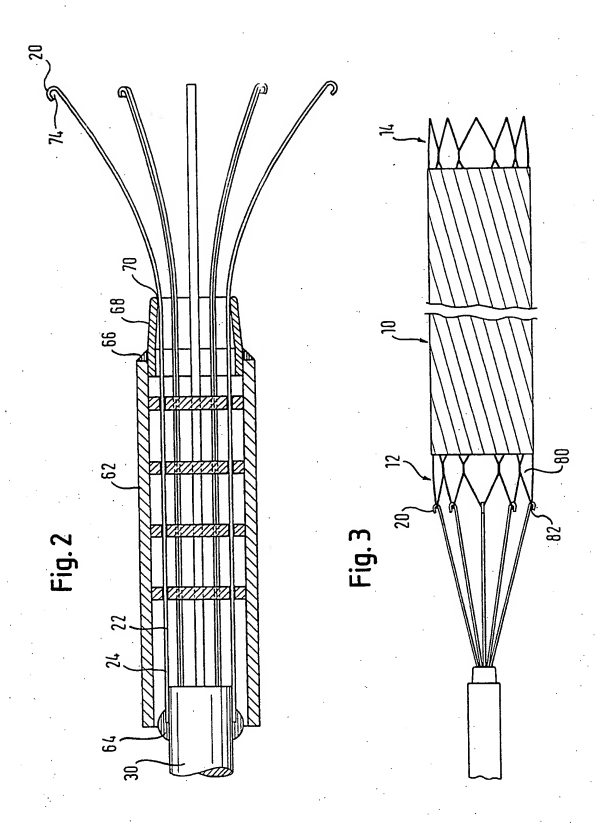
after drawing the stent through the constrictor and into the sleeve by the pull wires, thereafter removing the pull wires through the second end of the stent by pulling on the retraction wire.

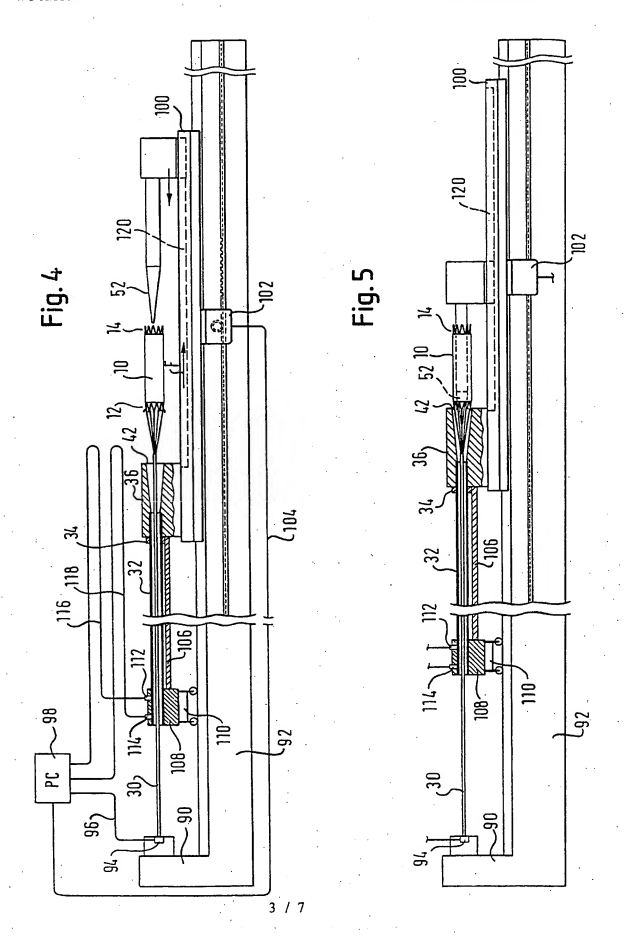
30 74. A method as defined in claim 70 further comprising:

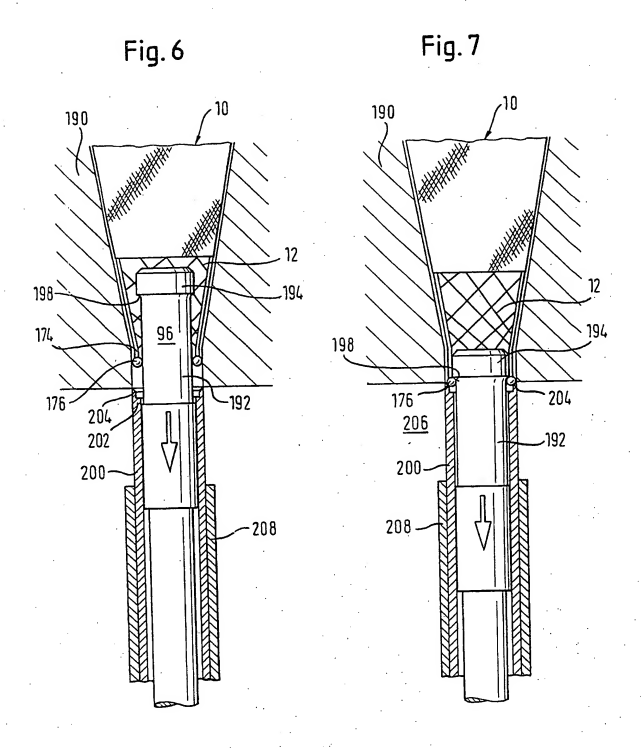
inserting a supporting tube into the lumen of the stent before drawing the stent through the constrictor.

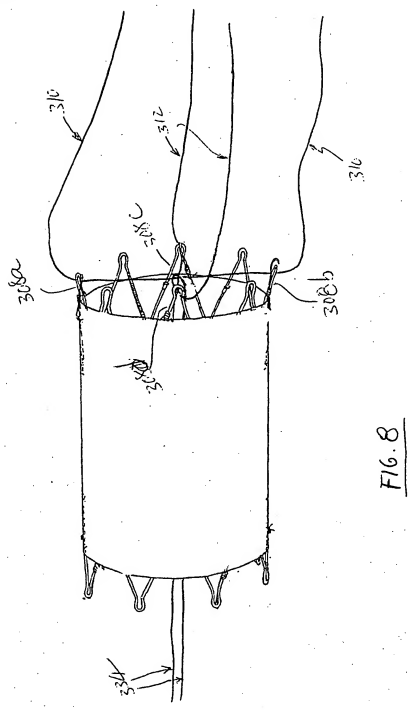


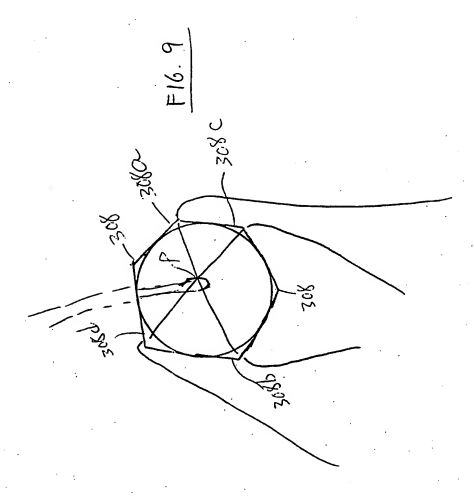
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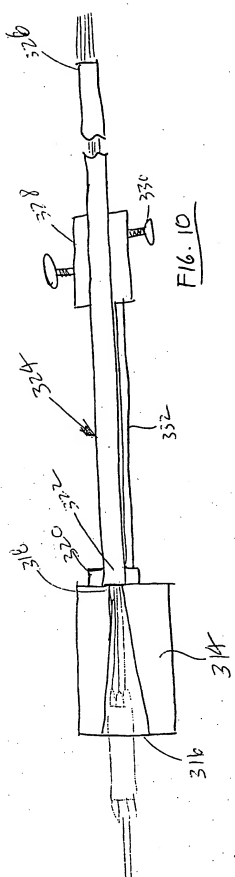


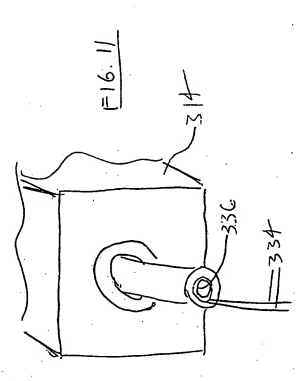












INTERNATIONAL SEARCH REPORT

Inten nai Application No PCT/US 00/29904

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 A61F2/06

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

 $\begin{array}{ll} \mbox{Minimum documentation searched (classification system followed by classification symbols)} \\ \mbox{IPC 7} & \mbox{A61F} \end{array}$

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal

Category *	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claim No.	
(WO 98 27894 A (PROGRAFT MEDICAL 2 July 1998 (1998-07-02)	INC)	1,50	
Ą	figures 8D-8F claims 44-47 page 24, line 17 -page 25, line	26	33, 64-68,70	
X	US 5 902 334 A (WEISER MICHAEL 11 May 1999 (1999-05-11) figure 15	ET AL)	50	
A .	column 11, line 54 -column 12,	line 43	1,33, 64-68,70	
· ·		-/		
X Fu	ther documents are listed in the continuation of box C.	X Patent family members are listed	in annex.	
"A" docum cons "E" earlie filing	categories of cited documents: ment defining the general state of the art which is not idered to be of particular relevance r document but published on or after the international date ment which may throw doubts on priority claim(s) or	 "T" later document published after the interpretation or priority date and not in conflict with cited to understand the principle or the invention "X" document of particular relevance; the cannot be considered novel or cannot involve an inventive step when the document. 	the application but early underlying the claimed invention to be considered to	
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